APPENDIX II

CLAIM No.	Serial No. 08/877,585
73. A coated optical fiber comprising: an optical fiber; and	Original claim 91 (page 150) is directed to "A coated optical fiber." Page 1, lines 17-19 ("The invention also relates to coated optical glass fibers")
a cured primary coating on the optical fiber	Original claim 103 (page 153) recites "said inner primary coating is formulated from a radiation curable coating." The cured primary coating is disclosed at page 34, lines 1-4 ("compositions were suitably cured by exposure to UV light"); page 49, lines 34-36 ("cured by exposure to UV light to form cured coatings.")
wherein the primary coating is strippable from a portion of the optical fiber	Original claim 104 (page 153) describes a primary coating that is strippable from a portion of the optical fiber, where strippability is described in terms of fiber friction Strippability of the primary coating is described on page 8, lines 19-33 (" said inner primary coating is adapted to provide a fiber friction force to allow the inner primary coating to slide readily off from the optical glass fiber when a stripping force is applied ")
at a temperature in at least a portion of the temperature range from about 25° to about 125° C	The temperature is described on page 34, lines 2-3 ("The dL/L for each coating was measured over the temperature range of 25 C (ambient temperature) to 125 C (highest usual stripping temperature).")
by exerting a force to a portion of the primary coating about the portion of optical fiber in a direction parallel to the longitudinal axis of the glass fiber which is away from a portion of the primary coating remaining on the optical fiber, such that the exertion of force,	A stripping method is disclosed on page 17, lines 32-39 ("When a typical ribbon stripping tool is applied to a ribbon assembly, pressure is applied to the ribbon assembly between heated plates. At the ends of the plates near the cut made in the matrix material and the inner and outer primary coatings, the inner primary coating can form an initial delamination site on the optical glass fiber, shown at 27 and 28 [in Fig. 1].")
followed by an optional one wipe with an alcohol laden piece of cloth or	An optional one wipe with alcohol is

CLAIM No.	Serial No. 08/877,585
paper of the stripped portion of the optical fiber, results in a stripped portion of optical fiber having substantially no residue of the primary coating.	disclosed in the article by Mills (the "Mills test") which is incorporated by reference at page 6, lines 27-32. The results of stripping is disclosed on page 8, lines 29-31 ("while leaving substantially no residue on the surface of said optical glass fiber during ribbon stripping"); page 50, lines 20-23 ("Microscopic examination of the pulled-out fibers at low magnification (e.g., 10X) clearly revealed the presence or absence of debris on the glass surface. If debris was present, the amount of debris was noted").
74. The coated optical fiber of claim 73,	
wherein the primary coating is strippable at the temperature upon execution of the force,	The disclosure of stripping at the temperature upon execution of the force is explicit throughout the disclosure and is mentioned at page 16, lines 14-21 in explanation of the fiber friction forces at stripping temperature.
to leave a stripped portion of the optical fiber having a Mill's value of about 2 or less.	The Mills' test values represent strip cleanliness, (page 58, lines 17-20: "When referring to strip cleanliness and predicted strip cleanliness herein, the numerical values correspond to those of the Mill's test."). Values of about 2 or less are described on page 110, lines 38.
75. The coated optical fiber of claim 73,	
wherein the primary coating is strippable at the temperature upon exertion of the force to leave	The disclosure of stripping at the temperature upon execution of the force is explicit throughout the disclosure and is mentioned at page 16, lines 14-21 in explanation of the fiber friction forces at stripping temperature.
a stripped portion of the optical fiber having a Mill's test value of 1.5.	A Mills' test value of 1.5 is described on page 60, Table 5, Coating F, Rating (Mill's Value 1.5) and page 84, Table 8 (Ex. 4-5, 4-7, 4-11, 4-12 all values were 1.5).
76. The coated optical fiber of claim 73, wherein the coating prior to curing is liquid at	The coatings prior to curing are liquid at 60 °C, as shown by the measurement of viscosity at 25 °C, as shown on page 74, Table 6,
are contains prior to curing is riquid at	(viscosity of primary coating compositions at

CLAIM No.	Serial No. 08/877,585
60° C.	25 °C, in the range of 6760 to 7650 mPa.s; similar viscosity values at 25 °C for the primary coating compositions in Tables 7, 8, 9 and 10).
77. The coated optical fiber of claim 76, wherein the coating prior to curing is liquid at 25 °C.	The coatings prior to curing are liquid at 25 °C, as shown by the measurement of viscosity at 25 °C: page 74, Table 6, (viscosity of primary coating compositions at 25 °C, in the range of 6760 to 7650 mPa.s; similar viscosity values at 25 °C for the primary coating compositions in Tables 7, 8, 9 and 10).
78. The coated optical fiber of claim 73,	
wherein the primary coating is made of a material which is removable from a glass substrate by an adhesion test force of less than 45 g/in, as measured by a peel back test at 50% relative humidity,	The adhesion force (at 50% RH) of 45 g/in is shown in Ex. 2-2, Table 3, page 53; and an adhesion test force of less than 45 g/in. is shown in Ex. 2-3, Table 3, page 53.
has an elongation to break of at least about 88% as measure by ASTM D-638, and	An elongation to break of at least about 88% is shown by the examples: Table 8, page 83, Elongation: 100% to 180% (Ex. 4-8, 4-9 and 4-10); Table 9, page 90, Elongation: 88% (Ex. 5-2).
has a tensile strength of at least about 72.5 psi as measured by ASTM D-638.	A tensile strength of 72.5 psi is disclosed in Ex. 5-2, Table 9, page 90 (0.5 MPa = 72.5 psi); Tensile strengths greater than 72.5 are disclosed for the primary coatings from Table 8, page 83: Ex. 4-8 (1.5 MPa = 218 psi); Ex. 4-9 (0.6 MPa = 87 MPa); Ex. 4-10 (1.1 MPa = 160 psi).
79. The coated optical fiber of claim 78,	
wherein the adhesion test force is about 14 g/in.	An adhesion test force of 14 g/in is disclosed in Table 3, page 53, for Ex. 2-3: Adhesion at 50% RH = 14 g/in.
80. The coated optical fiber of claim 73,	
wherein the primary coating is made of a material which is removable from a glass substrate by an adhesion force of at least 5 g/in, as measured by a peel back test at 95%	Actual reported values wet adhesion (95% RH) are disclosed in Table 3, page 53, adhesion force = 12 or 34 g/in at 95% RH (Ex. 2-3 and Ex. 2-2);

CLAIM No.	Serial No. 08/877,585
relative humidity, has an elongation to break is a least about 88% as measured by ASTM D-638, and	Elongation at break (measured by ASTM D-638M) of 88% is disclosed in Table 9, page 90, Ex. 5-2: Elongation = 88%; higher values are disclosed for examples Ex. 4-8 to 4-10 in Table 8, page 83.
has a tensile strength of at least about 72.5 psi as measured by ASTM D-638.	A tensile strength of at least about 72.5 psi is shown in Table 9, page 90, Ex. 5-2: (Tensile Strength = 0.5 MPa = 72.5 psi). Higher tensile strength values are shown in other examples: Ex. 5-4 (Tensile Strength = 1.1 MPa = 159.5 psi); Ex. 4-10 (Tensile Strength = 1.5 MPa = 217.5 psi) (Table 8, page 83).
81. The coated optical fiber of claim 78, wherein the elongation to break is about 140%.	The elongation to break of 140% is disclosed in Table 8 (cont), page 83; Ex. 4-9.
82. The coated optical fiber of claim 78, wherein the tensile strength is at least about 145 psi.	A tensile strength of 145 psi is disclosed in Example 4-1 in Table 7, page 80 (1 MPa = 145 psi). Higher tensile strength values are disclosed in Examples 4-8 (1.5 MPa = 217.5 psi) and 4-10 (1.1 MPa = 159.5 psi) (Table 8, page 83).
83. The coated optical fiber of claim 73,	
wherein said primary coating layer comprises a cured reaction product of from about 5 to	A primary coating layer which is the cured reaction product of from about 5 to about 80 wt.% of oligomers is disclosed on page 103, lines 4-5.
about 80 percent by weight of one or more acrylate- or methacrylate- terminated urethane oligomers.	Acrylate- or methacrylate terminated urethane oligomers are disclosed on page 102, lines 11-26 ("oligomers according to the invention can be made, for example, by reacting a diisocyanate compound with a long chain alkyl containing a hydroxy functional group and a radiation-curable functional group The urethane linking group is formed by the reaction of the isocyanate group with a hydroxyl group. In a second reaction the remaining isocyanate group is reacted with the as yet unreacted hydroxyl group of the compound Suitable examples of reactants containing a radiation-curable functional group and a hydroxy group are hydroxyethylacrylate or 2-hydroxypropylacrylate.").

CLAIM No.	Serial No. 08/877,585
84. The coated optical fiber of claim 83,	
wherein said primary coating layer comprises	Disclosure of a primary coating layer
a cured reaction product of,	comprising a cured reaction product of from about 0.1 to about 20% by weight of
in addition to the acrylate or methacrylate- terminated oligomer	additional monofunctional monomers is found on page 79, lines 9-10 ("Suitable amounts of mono-functionalized monomers have been
from about 0.1 to about 20 percent by weight of	found to be about 0.1 to about 20% by weight.")
a monomer selected from the group consisting of isobornyl acrylate,	Isobornyl acrylate is disclosed on page 115, line 27.
isodecyl acrylate,	Isodecyl acrylate is disclosed on page 90, Table 9 (Ex. 5-1, Ex. 5-3, Ex. 5-4).
hexanediol diacrylate,	Hexanediol diacrylate is disclosed on page 114, line 27.
phenoxyethyl acrylate, and	Phenoxyethyl acrylate is disclosed on page 97, Table 11, Ex. 10-22.
lauryl acrylate.	Lauryl acrylate is disclosed on page 115, line 28.
85. The coated optical fiber of claim 83 wherein said primary coating layer additionally comprises about 1.0 wt% of an organofunctional silane adhesion promoter.	The disclosure of organofunctional silane adhesion promoter is found on page 38, line 23 through page 40, line 23: ("'glass coupling moiety' is understood to mean a functional group which has the ability to improve adhesion to an inorganic surface or at an inorganic surface", "Silane coupling moieties are preferred."
	The amount of 1.0 wt.% is disclosed on page 74, Table 6, gamma-mercaptopropyl trimethoxy silane (1 part by weight of 100 parts).
86. The coated optical fiber of claim 85, wherein said adhesion promoter is a mercapto-functional silanes.	The mercapto-functional silanes are disclosed on page 40, lines 24-26 ("Common organic functionalities of the silane agents include, for example, mercapto,").
87. The coated optical fiber of claim 85, wherein said adhesion promoter is 3-	3-mercaptopropyltrimethoxysilane is disclosed on page 74, Table 6 (Ex. 3-1 to 3-4).

CLAIM No.	Serial No. 08/877,585
mercaptopropyltrimethoxy-silane.	
88. The coated optical fiber of claim 83, wherein said primary coating layer additionally comprises a photoinitiator.	The addition of a photoinitiator is disclosed on page 47, line 29 ("at least one photoinitiator").
89. The coated optical fiber of claim 83, wherein said primary coating layer additionally comprises from about 0.5 percent by weight of thiodiethylene bis(3,5-di-tert-butyl-4-hydroxy cinnamate.	A disclosure of about 0.5 percent by weight of thiodiethylene bis(3,5-di-tert-butyl-4-hydroxy cinnamate is in Ex. 2-4 on page 55, Table 4 (0.5%).
90. The coated optical fiber of claim 73, wherein the primary coating comprises the radiation-cured reaction product of the following ingredients:	A disclosure of "a radiation-curable inner primary coating composition containing at least one radiation-curable urethane oligomer comprising at least one polymeric block and at least one functional group capable of polymerization in the presence of actinic radiation connected to said at least one polymeric block" is found on page 11, lines 19-25.
(A) from about 5 percent to about 80 percent by weight of a reactively terminated urethane oligomer which is the reaction product of (i) polyether polyol; (ii) a wholly aliphatic polyisocyanate; and (iii) an endcapping monomer supplying a reactive terminus;	The amount of linear oligomer can be from about 5 to about 80 wt.% (page 103, lines 4-5). A disclosure of reactively terminated urethane oligomers is found on page 44, lines 23-30: "Representative linking compounds include diisocyanate compounds, wherein linkage occurs by formation of urethane, by reaction of hydroxyl, with isocyanate although aliphatic diisocyanates are preferred." Endcapping monomers supplying a reactive terminus are shown by the formula on page 98, lines 24 et seq., as linear oligomers to improve ribbon stripping operations: R¹-L-[R²-L] _n -R³, where R¹ and R³ are organic groupings having radiation curable functional groups; R² is an organic radical, L is a linking group, and [R²-L] _n can contain polyether (page 90, line 28).

CLAIM No.	Serial No. 08/877,585
	See also page 101, lines 8 et seq. for the radiation-curable oligomer of formula R ⁴ -x-L-x-[R ⁵ -x-L-x] _n -R ⁶ , where R ⁴ is a substantially linear long chain alkyl terminating in at least one hydroxyl group; L represent a molecular bridging group preferably derived from a diisocyanate precursor, x represent resulting reacting linking group, e.g., urethane, R ⁵ is a linear or branched or cyclic hydrocarbon or polyether moiety derived from a diol, and can contain a branched or cyclic aliphatic group, R ⁶ is an end group carrying a radiation-curable functional group.
(D) formalised 15 section 1 section 1	See also, e.g., page 112, lines 8-11 ("The radiation-curable oligomer can be easily formed by reacting a polymeric polyol, a compound containing a radiation-curable functional group and a hydroxyl group, and a polyisocyanate."); page 112, lines 16-19 (examples of polymeric polyols include polyether diols); page 113, lines 33 et seq. for organic polyisocyanates, including the wholly aliphatic polyisocyanates and examples in Table 15, page 117 and Table 16, page 118, Table 17, page 121, Table 18, page 123, Table 19, page 125, Table 20, page 127 and Table 21, page 129.
(B) from about 15 percent to about 65 percent by weight of ethoxylated nonyl phenol acrylate;	A disclosure of the inner primary coating composition including (B) ethoxylated nonylphenol acrylate between about 15% and about 65% is found in Ex. 11-1, Table 11, page 97 (15.1%) and Ex. 3-1, Table 6, page 74 (64.4%).
(C) from about 1.5 percent to about 3 percent by weight of a photoinitiator; and	From about 1.5 to about 3% of photoinitiator (C) photoinitiator is found in Table 8, page 75, Ex. 4-11 (1%), Ex. 4-10 (3%).
(D) about 1 percent by weight of an organofunctional silane adhesion promoter which binds in with the primary coating composition during cure;	about 1% of (D) organofunctional silane adhesion promoter is disclosed in Examples 3-1 to 3-4 in Table 6, page 74.
wherein all of the stated percentages are percentages by weight based on the total weight of the primary coating prior to cure,	The disclosure of the sums of the amounts based on the total weight of the primary coating prior to cure is seen from the data in the Tables wherein the total of the ingredients

CLAIM No.	Serial No. 08/877,585
	was 100 wt.%.
wherein the tensile modulus of the coating composition, when cured, is less than about 220 psi at 25° C, and	A disclosure of tensile modulus values of less than about 220 psi at 25°C is found in Table 8, page 83: Ex. 4-5 (1.2 MPa = 174 psi); Ex. 4-10 (1.1 MPa = 160 psi) and Ex. 4-8 (1.5 MPa = 217.5 psi).
wherein the refractive index of the cured coating composition is suitable for an optical fiber coating.	A disclosure of refractive index suitable for an optical coating is inherent throughout the specification. Formation of "clear" films suitable for an optical fiber coating are disclosed in Table 6, page 74.
91. The coated optical fiber of claim 73,	
wherein the primary coating comprising the radiation-cured reaction product of the following ingredients:	See discussion for Claim 90 for the support for claim 91.
(1) from about 5 to about 80 percent by weight of an acrylate- terminated aliphatic polyether urethane oligomers;	
(2) from about 15 to about 65 percent by weight of ethoxylated nonyl phenol acrylate;	
(5) from about 1.5 to about 3 percent by weight of a photoinitiator; and	
(6) about 1 percent by weight of an organofunctional silane adhesion promoter which binds in with the primary coating composition during cure;	
wherein all of said percentages being percentages by weight based on the weight of the primary coating prior to cure.	
92. The coated optical fiber of claim 91,	The coatings prior to curing are liquid at 25
wherein said primary coating is obtained by curing a composition that is liquid at 25° C.	°C, as shown by the measurement of viscosity at 25 °C: page 74, Table 6, (viscosity of primary coating compositions at 25 °C, in the range of 6760 to 7650 mPa.s; similar viscosity values at 25 °C for the primary coating compositions in Tables 7, 8, 9 and

CLAIM No.	Serial No. 08/877,585
	10).
93. An optical ribbon comprising a plurality of optical fibers of claim 73 and a matrix material, the plurality of fibers held together in a parallel arrangement by the matrix material.	A ribbon assembly comprising a plurality of coated optical fibers, at least one optical glass fiber coated with at least an inner primary coating and a matrix material bonding the plurality of coated optical fibers together is the subject matter of original claim 1. A disclosure of ribbon assemblies is found on page 1, lines 19-21 and page 131, lines 1 et
	seq.
94. A method of preparing a coated optical fiber for splicing,	A disclosure of a method for preparing a coated optical fiber for splicing including ribbon stripping is found on page 3, lines 3-
the coated optical fiber being a coated optical fiber according to claim 73,	20.
comprising the steps of:	
stripping the primary coating away from a portion of the optical fiber at a temperature in at least a portion of the temperature range from about 25° to about 125° C. by cutting with a blade into the primary coating,	The temperature of from 25 °C to 125 °C is disclosed on page 34, lines 2-3.
then having the blade exert a force on the primary coating in a direction parallel to the optical fiber to force the primary coating away from the portion of the optical fiber,	
and optionally wiping the portion of optical fiber from which primary coating has been forced away with an alcohol laden piece of cloth or paper,	A disclosure of the optional wiping step is found in the Mills' test incorporated by reference into the application on page 6, lines 27-32.
such that the stripped portion exhibits little or no residue of the primary coating.	The result of stripping is disclosed on page 8, lines 29-31 ("while leaving substantially no residue on the surface of said optical glass fiber during ribbon stripping"); page 50, lines 20-23 ("Microscopic examination of the pulled-out fibers at low magnification (e.g., 10X) clearly revealed the presence or absence of debris on the glass surface. If debris was present, the amount of debris was noted").

CLAIM No.	Serial No. 08/877,585
95. A method of splicing coated optical fibers comprising, preparing two optical fibers for splicing according to the method of claim 94; aligning the stripped portions of the prepared optical fibers; and joining the stripped portions of the prepared optical fibers.	A method for connecting optical fibers individually or as ribbon assemblies for "mass fusion splicing operation" is disclosed on page 2, lines 24-34.
96. A spliced optic fiber made by the method of claim 95.	A disclosure of a spliced optic fiber by the splicing operation is found on page 2, lines 24-34.
97. A coated optical fiber comprising: an optical fiber;	A coated optical fiber comprising an optical fiber and primary coating is disclosed on page 9, lines 8-12;
a primary coating coated onto the optical fiber, the primary coating made of a material which is removable from a glass substrate by an adhesion test force of less than 45 g/in, as measured by a peel back test at 50% relative humidity	Examples 2-2 and 2-3 (Table 3, page 53) show that the invention encompasses coatings having an adhesion force at 50% RH of 45 g/in or less.
an elongation of at least about 88% as measured by ASTM D-638, and	An 88% elongation is disclosed in Example 5-2 on page 90 in Table 9; ASTM D-638M is disclosed on page 63, line 8 to page 65, line 18.
a tensile strength of at least about 72.5 psi as measured by ASTM D-638.	A tensile strength of 72.5 psi, is disclosed as 0.5 MPa in Example 5-2 on page 90 in Table 9. The conversion from MPa to psi (1MPa = 145 psi) can be found on page 64, line 5.
98. A process for preparing a coating optical fiber comprising (1) applying to an optical fiber a primary coating composition layer comprising a mixture of the following ingredients:	A process for preparing a coated optical fiber by applying a primary coating composition to the optical fiber is shown generally throughout the specification and examples, including, page 1, lines 24-26; page 48, lines 17-29 (including the disclosures of incorporated patents 4,474,830 and 4,913,859).
(A) from about 5 percent to about 80 percent by weight of a reactively terminated urethane oligomer which is the reaction product of (i) a polyether polyol; (ii) a wholly aliphatic polyisocyanate; and (iii) an endcapping	Compositions with ingredients (A), (B), (C) and (D) are also described throughout the specification. A reactively terminated urethane oligomer (see, e.g., the formulas on page 45, lines 8-19; page 98, lines 25-38;

Serial No. 08/877,585
page 101, lines 10-22) wherein the linking group L or x is preferably urethane (e.g., page 45, line 22; page 98, line 35; page 101, line 21).
The reaction of a mixture of (i) polyether polyol, (ii) wholly aliphatic diisocyanate and (iii) end-capping monomer supplying a reactive terminus, is shown in Examples (e.g., page 53, Table 3, Oligomer C: reaction product of (i) polytetramethylene ether glycol, (ii) isophorone diisocyanate, and (iii) hydroxyethylacrylate and mercaptosilane) and is also generally disclosed on page 112, lines 8-11 ("The radiation-curable oligomer can be easily formed by reacting a polymeric polyol, a compound containing a radiation-curable functional group and a hydroxyl group, and a polyisocyanate."). The range of about 5 to about 80% is disclosed on page 94, lines 1-6.
The Examples on page 53, Table 3, also include (B) ethoxylated nonyl phenol acrylate; (C) photoinitiator; and (D) organofunctional silane, at or within the stated ranges. The range of about 15% to 65% for reactant (B) is found in the examples, e.g., page 97, Table 11; page 74, Table 6, respectively. The range of about 1.5% to about 3% for reactant (C) is found in the examples, e.g., page 32, Table 1 (Comp. Ex. A-2 1.54%); page 97, Table 11 (3%). The amount of the organosilane adhesion promoter is 1% in the composition of
Example 10-22, page 97, Table 11.
The tensile modulus of the cured composition of less that about 220 psi is shown to be part of the invention by reference to, for example, Example 4-1, on page 80, Table 7 (1.4 MPa =
203 psi); Ex. 4-5, Ex. 4-9 and Ex. 4-10, on
page 76, Table 8 (CONT) with values of 174
psi (1.2 MPa), 160 psi (1.1 MPa) and 188.5 psi (1.3 MPa), respectively.
The refractive index being suitable for an
optical coating, is implicit throughout the disclosure and is exemplified by the clear

CLAIM No.

suitable for an optical fiber coating;

and

(2) radiation-curing said coating in situ,

wherein the primary coating is strippable from a portion of the optical fiber at a temperature in at least a portion of the temperature range from about 25° to about 125° C. by exerting a force to a portion of the primary coating in a direction parallel to the longitudinal axis of the glass fiber which is away from a portion of the primary coating remaining on the optical fiber, such that the exertion of force followed by an optional one wipe, with an alcohol laded piece of cloth or paper of the stripped portion of optical fiber, results in the stripped portion of optical fiber having a substantially no residue of the primary coating.

Serial No. 08/877,585

films obtained in Examples 3-1 to 3-4, on page 74, Table 6.

The step of radiation-curing in situ is also implicit throughout the disclosure and is shown, for example, on page 34, lines 1-2 and page 49, lines 34-36.

That the primary coating is strippable at the temperature ranges and under the conditions specified, with substantially no residue of the primary coating is disclosed throughout the specification, for example, page 34, lines 2-4 for the temperature range of 25° to 125° C; and Ex. 2-1, on page 51, Table 2, for "no residue." The optional one wipe with alcohol is disclosed in the incorporated Mills test disclosure as stated on page 6, lines 27-32.